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HRnD to evitevise alregitnA (43)

(57) The invention concerns a conjugate of the formula: Pyr-His-Trp-Ser-Tyr-D, Lys-Leu-Arg-Pro-Y,

89 (cc)

Pyr = pyroglutamic acid :nieterlw

Trp = tryptophan enibitain = aiH

Tyr = tyrosine euues = 1eS

ren = jenciue D.Lys = D. Lysine

= broune ᅃ = sudiuius DIA.

= GIY NH2 or NHEI

An immunogenic substance capable of raising antibodies to GnRH in a mammalian subject, and which comprises the Pro-Y as defined above. ■ an immunogenic carrier protein preferably diptheria toxoid or tetanus toxoid, or Pyr-His-Tro-Ser-Tyr-D.Lys-Leu-Arg-

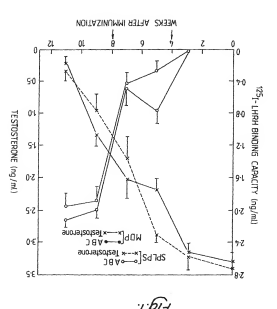
domestic pets, the treatment of breast cancer, endometrosis, precocious puberty, the treatment of cancer of the prostate antagonist of Griffit (LHRH) may be usefully used, e.g. the control of male and female fertility, the suppression of heat in Since GriRH is a control hormone, the conjugate and/or immunogenic substance is useful in all situations where an above conjugate is also provided.

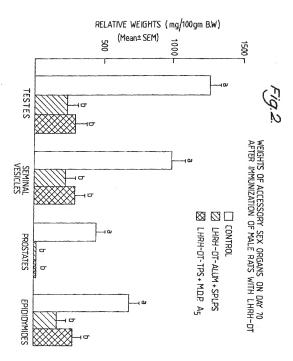
and as a post-partum contraceptive.

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This print takes account of replacement documents submitted after the date of thing to enable the application to comply with the formal At least one drawing originally filed was informal and the print reproduced here is taken from a later filed formal copy

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Antigenic Derivative of GRRH

The present invention relates in general to the control of fertility and the treatment of fertility associated conditions. More particularly (but not exclusively so) it relates to carcinoma of the prostate in males and to a process for the preparation of an improved anti-chRH vaccine, which on application of an subjects causes atrophy of the prostate and thereby subjects causes atrophy of the prostate and thereby autentially diminishes the area within which carcinoma

It is well known that carcinoma of the prostate is a wide-spread syndrome in males and in a large percentage of cases, its occurrence and growth are directly hormones are produced in the testes, doctors have in past resorted to orchiectomy, i.e. operation for removal of the testes, in order to do away with the source of the testes, in order to do away with the source of the testes, in order to do away with the source of the testes, in order to do away with the source of the testes.

It is also known that the decapeptide, gonadotropin release hormone (GRRH also referred to as LHRH), which is present in the body regulates male sex hormone production in the testes by virtue of its stimulatory action on the in the testes by virtue of its stimulatory action on the fin the testes by virtue of its stimulatory action on the in the production. A direct stimulatory causing release of gonadotropins. A direct stimulatory causing representation of programmer is not constant.

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can occur.

excluded.

Therapeutic utility of superactive analogues of GRRH

286:1607-1609, 1983). Drawbacks are the high cost of Jonadotropin releasing hormone agonist. Br. Med. J. yqxsuceq csrcruows of the brostste: Trestment with s 52 Williams G, Bloom SR: JM, O'Shea JP, Mashiter K, prostate cancer. Br. Med. J. 286:1309-1312, 1983. Allen with gonadotropin releasing hormone analogue in advanced Whititeld HM, Besser GM, Malpas JS, Oliver RTD: Treatment prostatic carcinoma (Waxman JH, Wass JAH, Hendry WF, 20 (IHRH) have proved useful in the treatment of advanced Several potent agonist analogues of GnRH curourcettă. rednistion, which occurs when analogues are administered phenomenon of pituitary desensitization or down-1981, pp 321-333). Such applications are based on the SI Contraceptives". Philadelphia: Harper & Row Publishers, JD, Scierra JJ (eds): "LHRH Peptides as Female and Male hypogonadotropic hypogonadism. In Zatuchni GI, Shelton WF, Vale WW, Rivier J, MacArthur JW: LHRH in Proc. Natl. Acad. Sci. USA 79:1658-1662, 1982. Crowley OT with luteinizing hormone releasing hormone agonists. tubibition in patients with prostatic carcinoma treated ATA, Camaru-Schally AM, Schally AV: Tumor growth Tolis G, Ackman D, Stellos A, Mehta A, Labrie F, Fazekas Proc. Soc. Exp. Biol. Med 175:259-281, 1984. susjodnes of hypothalamic hormones in endocrine-dependent Comexu-Schally AM, Redding TW: Anti-tumour effects of (Schally AV, abnormalities has been demonstrated. (LHRH) to ameliorate a spectrum of androgen dependent

releasing hormone from injectible microcapsules. Proc courtoffed refesse of [D-fip]6 internizing hormone skatems tor peptides: inhibition of rat prostate by Schally AV, Tice TR, William EM: Long acting delivery cexcinome, the Lancet 2:1201-1205, 1985. Redding TW, scting D-TRP-6-LHRH microcapsules in advanced prostate Mandomised controlled study of orchidectomy vs long-SL, Allen L, Phillips RH, Edwards L, Schally AV: at which they must be administered. (Parmer H, Lightman tyese combonugs sug' except tu s tew cases the treduency

Watl. Acad. Sci. USA 81:5845-5848, 1984).

treatment with a stimulatory LRH analogue - a new C, Wide L: Inhibition of ovulation in women by chronic York: Reven Press 1984 pp351-359. Willius SJ, Berguist "Hormone receptors in growth and reproduction". In Saxena BB, Catt KJ, Birmbauma L, Maritini L, (eds): immunization against gonadotropin releasing the hormone. pormone: potential uses of active and passive bitnitery sites of action of gonadotropin-releasing Study O, Das C, Gupta SK, Singh G: Pituitary and extra produced by GRRH (LHRH) agonists. (Talwar GP, Singh V, primates have sex-steroid profiles similar to those inhibitory effect on the pituitary-gonad axis. Immunized with the hormone. Active immunization leads to an surfacebased by antibodies that are specifically reactive The biological activity of GRRH (LHRH) can also be

approach to birth control. Contraception 17:537-545,

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sudeuger suff-Gugh (IHRH) response with human compatible was conjugated to tetanus toxoid [TT], and which could was employed. An alternate modality in which GnRH (LHRH) combjete adjuvant, which is nonpermissible for human use, 55:616-622, 1973.), but in these studies Freund's tropin releasing hormone, Biochem. Biophys. Res. Commun. characterization of an antiserum to synthetic gonado Erigkin M, Chobsung P, Zor V, Lindner HR: Production and коси д' мттсрек м' *\$46T '90\$-6EE:E9 Endocrinol. testes and accessory sex organs in the male rat, releasing hormone on serum and pituitary gonadotropins, Effect of scrive immunization to inteinizing hormone 1103, 1973. Fraser HM, Gunn A, Jeffcoate SL, Holland DT: of radioimmunosessy for LHRH, Endocrinology 93: 1092associated with gonadal atrophy in rabbits. Development production of antiserum to LH releasing hormone (LHRH) Kumasaka T, Worobee RB, Dunn L, Debeljuk L, Schally AV: previously by several investigators, (Arimura A, Sato H, 1978). Bioeffective immune response has been generated

antibody response without Freund's complete adjuvant against a "self" peptide, luteinizing hormone releasing hormone (LHHH), Am. J. Reprod Immunol 1:262-265, 1981).

EP 1812546 Firman-Moore Inc., discless the use of

adjuvants, has been described (Shastri W, Manhar SK, Talwar GP: Important role of the carrier in induction of

EP 181236A2 Fitman-Woore Inc., discloses the use of conjugates comprising anologues of GnRH which can be used as an anti-LHRH vaccine to prevent the function of LHRH

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This application suffpodies active against LHRH. THEH MUTCH CON DO need so an immunoden to broduce qtacjoses tye nae ot conjudates compristing analogues of UK 2,196,969A Proteus Biotechnology Ltd., similarly

US 4,676,981 D.W. Silversides et al. discloses the applicability to prostate cancer. mentions that the vaccine disclosed may have

bassive immunisation of these antibodies affects sex in vitro production of antibodies to GnRH, and that

stimulating hormone (FSH) analogue together with a ST ot a luteinizing hormone (LH) analogue or a follicle Ascertasry vaccine comprising a protein-hormone conjugate WO 88/01176 M.R. Brandon discloses a contraceptive gland weight.

the present invention. However, the conjugates disclosed by these documents protein hormone releasing hormone (LHRH) analogue.

the GnRH peptide analogue of through an amino acid immunogenic cerrier substance or to enother molecule of the GnRH peptide analogue is conjugated to either an tuvention and as distinct from the foregoing disclosures, Furthermore, in the present differ in respect of the analogues of GraH provided by

.eupolens jocated centrally within the peptide chain of the

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tor preparing GraH (LHRH) analogue conjugates of The present application describes an improved method

replacement of glycine at position 6 by D-lysine. consistent immunogenicity. The peptide backbone of GnRH

(LHRH) was modified to engender an amino group by

was optionally linked to t-amino caproic acid p-alanine

or other non-protein amino soid, which has a functional

droup for ensuring conjugation to an immunogenic carrier

protein or to another modified peptide backbone of GnRH

His = histidine wyereru byr = pyroglutemic acid

Pyr-His-Trp-Ser-Tyr-D-Lys-Leu-Arg-Pro-Y

ratsing antibodies to GnRH in a mammalian subject,

there is provided an immunogenic substance capable of yecciqtud to one sabect of the present invention

prostate. The vaccine is long lasting in its effect, and effect mey be block of fertility or an atrophy of the

level of male or female sex hormones. An accompanying this down regulation, there is a drastic reduction in the which down regulate the action of GnRH. As a result of effcits within the body the production of antibodies s Ascerne which when applied to a mammalian subject yeccordingly, the invention concerns the provision of

combitatud e coulndate of the tormnie:

does not require frequent medication.

Ltb = fxXbfobpsu

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· (HMHI)

Tyr = tyrosine Ser = serine

D-rat = D-lysine

ren = jencjue

Arg = arginine

Pro = proline

GTA = dTActue

-NHEC) * $x = -c_1 \lambda_{MH}^{5}$ or -Met (slso sometimes designated as

N = 90 rmmnuodeurc cerrter broteru

The conjugate may be accompanied by a suitable

adjuvant, optionally after adsorbing the conjugate on

The immunogenic carrier protein may be coupled to · mule

e.g. hydroxylysine, a-amino adipic acid, a-amino/8this purpose, other non-protein amino soids could be used swinoceproic acid and p-alanine are especially useful for SO are unusual non-protein amino acids. ~> ⊅stidw peptide and protein. E-aminocaproic acid and B-alanine slanine substituent to define the moler ratio between the sminoceproic soid (amino-hexanoic soid or AHA) or B-Preferably, the D-Lys residue is provided with a E-1(3-dimethyl-amino-propyl)-3-ethyl carbodiimide. the D-Lys residue using for example glutaraldehyde or

(ECDI) is a compling reagent which activates the carboxyl 1(3-qimethyl-smino-propyl)-3-ethyl carbodifmide conjugation is made through the NH2 grouping. ys In b-signine (H2N-CH2-CH2-COOH) sarcosine.

smino/a-dismino/ &-dismino butyric acid, ornithine or

sre linked thereby. Examples of the carrier protein include diphtheria

Blochem. (1976), 79, 233-236).

The immunogenic substance in the form of a dimer may glutaraldehyde wherein respective f-amino groups of the complete substance in the two peptides reing glutaraldehyde and glutaraldehyde son

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maleimidophyenyl) butyrate; Kitagawa, T. et. al., J. (1976), 79, 233-236), or SMPB (succinimidyl 4-(psuccinimide ester; Kitagawa, T. et. al., J. Blochem. (1979), 101, 395-399), MBS (m-Maleimido benzoyl-N-hydroxy carboxylate; Yoshitake, S, et. al., Eur.J.Biochem. (snccivimidyl 4-(N-maleimidomethyl) cyclohexane-1et. al., Biochem. J. (1978), 173, 723-737), SMCC succinimidyl 3-(2-pyridyl dilhio) propionate; Carlsson, J protein can be employed. For example by use of SPOP (Nconjugation to the NH2 or COOH group of the carrier functional -MH2 group several other methods of Hearud created a b-alanine to form the conjugate. with the 6-amino group of the AHA or the 6-amino group of is linked to D-Lys, the activated-COOH group will couple smino group of D-Lys or \$-alanine. If AHA or \$-alanine cerpoxit droup of the cerrier protein can attach to 6-The ECDI activated AHA or \$-alanine for conjugation. the conjugate. This coupling reagent does not require the amino group of the other peptide or protein to form drond of a peptide or a protein which in turn reacts with The structure of the peptide set out employs conventional abbreviations with the amino groups of each amino acid appearing to the left and the carboxyl groups to the right. The first five and the last four amino order as the amino acide of GnRH. At position 6, the glycine of GnRH has been replaced by D-lysine and it is through the 6-amino group of the D-lysine that, for example, 6-amino group of the D-lysine that, for example, 6-amino oppoic acid (amino-hexanoic acid or 6-amino group of caproic scid (sanno-hexanoic acid or 6-amino group of caproic scid is available for conjugation to the macromolecular protein carrier by the conjugation to the macromolecular protein carrier by the

Fig.2 shows a bar graph of weights of accessory sea organs on day 70 after immunisation of male rats with

GARH-DT.

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Fig.1 shows antigen binding capacity (ABC) and restorterone levels in rats immunized with GnRH-DT;

detail with reference to the drawings in which:

conjugate in the preparation of an anti-GARH vaccine. In order that the present invention is more fully understood embodiments will now be described in greater

The invention further includes the use of the above

The invertion also includes the above conjugate for $$\operatorname{Top}_{\mathbb{R}^n}(X)$$

toxoid (DT) and tetanus toxoid (TT). In the case of the dimer, the carrier protein Σ is provided by the other

the next protected amino acid coupled to the preceding escy confirma, the amino-protecting group was removed and were purchased from Bachem and Sigma Companies. After conbjeq ou. All protected amino acids used for synthesis MH2 group, protected amino acids were successively 50, 1981. Starting with the resin which possesses a free John M Stewart in their work "Peptide", Volume 2, PP 45secondance with what is described by Gary R Matsueda and This resin can be prepared in benzhydrylamine resin. tuvention employing as solid support para-methyl tormula was synthesised according to the present peptide synthesis, the peptide of the above-mentioned Based on the established methodology of solid phase Pierce Chemical Company, Rockford, Illinois, USA, 1984. their book entitled "Solid Phase Peptide Synthesis", peer descriped by John M Stewart and Janis D Young in beptide synthesis and the techniques involved have

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quitaraldehyde method.

With the exception of amino hexanoic soid and gyrine which have no chiral centre and lyaine which is of D-configuration, all the amino acids of the peptide of L-configuration. The choice of D-lyaine instead in the body of the subject than L-lyaine and secondly in the body of the subject than L-lyaine and secondly in the body of the subject than L-lyaine and secondly agonistic behaviour with respect to native GNRM:

bebrige was then conjugated to a carrier protein to Lye baritied rc3000 Skatem) natud s Akqsc C18 colnmu. the peptide was effected by preparative HPLC (Waters Prep de-protection of the protecting groups. Purification of aceaeuder which action also resulted in the simultaneous ridnig placeder tinoride with entsole present as a peptide was cleaved off the resin by means of anhydrous After the synthesis was complete, the .07e1 ,8e2 oa publication "Analytical Biochemistry", Volume 34, pp 595 R.L. Colescott, C.D. Bossinger and P.I. Cook in their in accordance with the method described by E. Kaiser, one. The coupling and de-protection steps were monitored

rert-putyloxy carbonyl (Boc) recognised abbreviations are as follows: SI

blucolintemic sciq (blr) (psuzyl (bzl) y-promopenskionk carponly (Brz) 20 peuzhjoxh-cerpouhj (z) 0-Ifnoxenyl methoxy carbonyl (Fmoc) b-togneue anggoulg (Los)

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Lys(NcBoc), Nc-Z-AHA, Boc-Tyr(Brz), Boc-Ser(OB3L), Bocpoc-dja, boc-pro, boc-Arg(Tos), boc-Leu, NuFmoc-Dconfige is as tollows:

The order in which the protected amino acids are

The amino-protecting groups employed with their browide the immunological regions of the desired vaccine.

Trp, Boc-His(Tos) and Z-Pyr.

on snalysis, AHA enables quantification of the number of to link the peptide to the carrier protein. Furthermore, WHY is su numens; swino soid the purpose of which is

The preferred coupling agent employed for the above moles of peptide which are linked to the protein.

Myere the amino-protecting group is Boc, removal mentioned step is dicyclohexyl carbodifmide (DCC).

sceric sciq in dichloromethane followed by neutralisation OT tyereof is preferably effected by means of 50% trifluoro

species to preferrebly effected by means of 20% piperidine Myere the amino-protecting group is Fmoc, removal with 10% triethyl smine in dichloromethane.

coupling of the first amino acid. This treatment is effected after each amino acid. protecting group is carried out after the coupling of The sequence of steps for removal of the amino-

trestment is effected for one minute (unless otherwise fAbrosy qe-brotection sednence in which each wash effected after coupling of the first amino acid. brotection seguence in which each wash treatment is w typical de-

Three-time wash with dichloromethane stated) is as follows:

scid in dichloromethane containing 1,2 Mash for five minutes with 50% trifluoro acetic

in dimethyl formamide.

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- 9. Wash for thirty minutes with 50% trifluoro acetic acid in dichloromethane containing 1%
- 1,2-ethane dithiol
- 4. Two time wash with dichloromethane
- 5. Two time wash with 1% 1,2 ethane dithiol in
- tsopropyl alcohol 6. Three time wash with dichloromethane
- γ_{\star} . Wash for two minutes with 10% triethylamine in
- qicyjoromethene
- 8. Wash for ten minutes with 10% triethylamine
- enedtemoroldoth ditw daew emit eerd? .
- 9. Three time wash with dichloromethane.

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- After each de-protection sequence is completed, the successive amino acid to be coupled is then added, in two-fold excess together with dicyclohexyl carbodismide as coupling agent. The coupling reaction
- proceeds for approximately two hours.

 With the exception of the instances identified hereafter, the solvent medium employed throughout the 20 coupling and de-protection reactions is dichloromethane.
- coupling and de-protection reactions is dichloromethane.
- When coupling N-Fmoc-D-Lys(N Boc) and removal of Fmoc, the solvent employed is dimethyl formamide;
- Finec, the solvent employed is a mixture of dimethyl formamide.

 25 the solvent employed is a mixture of dimethyl formamide
- yffer configure N-Fmcc-D-TAs(NPBoc), the NPBoc is and dichloromethene:

removed employing 50% trifiuoro acetic acid-

smine in dichloromethane followed by the subsequent dichloromethane mixture, neutralised with 10% triethyl

SO% piperidine in dimethyl formanide and coupled with After coupling of Z-AHA the N Fmoc is removed with coupling on Z-AHA.

After it has been synthesised, the peptide is given BOC-TYY(BIZ).

scavenger in a reaction time of approximately one hour at embjoying anhydrous hydrofluoric acid with anisole as a dichloromethane mixture, then with methanol before being a final wash with a 50% trifluoro acetic acid-

peptide-resin mixture is washed with ether. The peptide SI O.C. Volatiles present are removed under vacuum and the The peptide is then cleaved off the dry resin

the present invention and the effect of the vaccine on constituting the immunological agent of the vaccine of breparation of the peptide-carrier protein conjugate The purification of the extracted peptide, the

is extracted with 10% scetic soid and lyophilised.

treated subjects are described in detail in the following

burilication of the extracted peptide was effected Purification of the Extracted Peptide

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Examples.

natud s Maters Prep-Lc 3000 liquid chromatograph. pk reverse-phase high performance liquid chromatography

EXYMPLE 1

cartridge or column of the chromatograph was of polyethylene 30 x 5 cm ib, hand-packed with Vydac C_{18} having a particle size of from 15 to 20 µm. The purification was carried out in two steps each employing a buffer solution consisting of two solvents. The buffer adueous triethyl ammonium phosphate of pH 2.5 (A) and 60% acceptorization for the second aqueous triethyl ammonium phosphate of pH 2.5 (A) and 60% acceptorization for the second scetto said in water (A) and 60% acceptorization C_{18} trifiuoro sectio said in water (A) and C_{18} second acceptorization sieg consisted of squeous C_{18} trifiuoro sectio said in water (A) and C_{18} second acceptorization C_{18} consisted of squeous C_{18} trifiuoro sectio said in water (A) and C_{18} sectionization C_{18} consisted of squeous C_{18} trifiuoro sectio said in water (A) and C_{18} sectionization C_{18} consisted of squeous C_{18} trifiuoro sectio said in water (A) and C_{18} sectionization C_{18} trifiuoro sectio said in water (B) and C_{18} sectionization C_{18} trifiuoro sectio said in water (A) and C_{18} sectionization C_{18} trifiuoro sectio said in water (B) and C_{18} sectionization C_{18} trifiuoro sectio said in water C_{18} sectionization C_{18} trifiuoro section C_{18} sectionization C_{18} trifiuoro section C_{18} sectionization C_{18} trifiuoro section C_{18} trifiuoro section

The fractions resulting from the first purification step were collected in samples of approximately 75 ml each and isocratically analysed in aqueous acetonitrile containing 0.1% trifluoro acetic acid. Those fractions which resembled each other most and which appeared to be pure were pooled separately. Each pool was diluted to 1 litre by the addition of triethyl ammonium phosphate and reloaded into the chromatograph in separate runs for the second step of purification. Once again, the fraction resembling each other most and appearing to be pure were pooled separately and the pooled amounts lyophilised.

nm and chart speed 1 cm per minute.

Every 1.5 g of crude peptide subjected to putification by this two-step liquid chromatography yielded 650 mg of pure peptide. On analysis, it was found that the amino soid composition of the putified

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bebirge corresponded to its constituent amino acids as

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1.23	:	skī	1.24	:	yrd
1.14	٠.	ue-I	ε.1	:	stH
1.27	:	TYr	80.1	:	GJA
86.0	:	AHA	77.0	:	zes
0.τ	:	Pro	81.1	:	Pyr

EXYMBLE 2

4°C with three changes. "Spectrapor" (Trade Mark) 18 O.V Hq to noitules relited standard M I.O to sertil SI whereafter the reaction was stopped by dialysis against 97. for 20 hours in a mechanical shaker in a cold room, thereof in the mixture was 0.1%. The mixture was shaken after each addition. The concentration of glutsraldehyde bebirge-giburperis roxotq mixture which was shaken well coored in ice and slowly added at 5 ml a time to the are 45 ml of 0.7 Hq to entiter reline of M 1.0 to im 24 ni (Sigma grade II (Trade Mark), 25% w/v aqueous solution) mixture kept in cold condition. 234µl glutaraldehyde phosphate buffer saline of pH 7.0 were added and the M 1.0 to im 00 ni (9nne, pune) in 60 ml of 0.1 M St solution, 28.125 mg diphtheria toxoid (obtained from pH 7.0 and cooled in ice. To the cooled peptide to entire relited standardoff M I.O to Im Z nt beviosaib asw 40mg of the peptide prepared according to Example 1 Preparation of Peptide-Diphtheria Toxoid Conjugate OT

Immunochemistry 6: 43-47, 1969). conjugate for the detection of antigen and antibodies. use of the enzymes to proteins with glutaraldehyde. dinteraldehyde conjudation see Avrameas S: Coupling of (For further details on .besu asw 000,01 lo dialysis tubing having a molecular weight cut-off limit

column employing a 0.1 M sodium-phosphate buffer having a The conjugate was finally purified over a LKB TSK 3000 SW Mark) membrane filter having a cut-off limit of 30,000. concentrated by ultrafiltration using an "Amicon" (Trade After dialysis, the formed conjugate was

Preparation of Peptide-Tetanus Toxoid Conjugate EXYMPLE 3

purified peptide of Example 1 were employed and 37.5 mg Example 2 above with the exception that 36.56 mg of the The procedure followed was the same as that of

of telenus toxoid was substituted for the diphtheria

swino scid, amino caproic scid, which is present only in ph swruo sorg susphers taking squantage of the unusual i.e. diphtheria toxoid and tetanus toxoid, was estimated ot conjugation of the peptide to the carrier proteins, In respect of each of Examples 2 and 3, the degree toxotd.

coulndstion of the peptide was found to be from 10 to 25 The degree of the peptide and not in the protein.

mojes ber moje of the carrier protein.

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EXAMPLE 4

Immunisation of Subjects Employing Vaccine Containing the

Peptide-Protein Conjugate as immunological Agent
Outbred adult male rate bred from an initial Wister

5 strain were injected according to an injection schedule
consisting of three intra-muscular injections of the
conjugate of the present invention. The injections
comprising 20 up per rat were given on contralateral
sites at monthly intervals. Thereafter the animals were

10 bled at fortnightly intervals from the retro-ovbital

plexus and the sera was stored at -20°C until assayed.

One group of ten rate was immunised employing the conjugate adsorbed on alum with 0.1 mg sodium phthalylated derivative of salmonella enteritidis ascend group of ten rate received nor-Muramyl dipeptide group, all the ingredients were in aqueous phase. For the second group, a water-in-oil emulaion was necessary to for which a vehicle composed of Tween 80 (Trade Mark), and the month of the first the month of the first the second group, a water-in-oil emulaion was necessary to for which a vehicle composed of Tween 80 (Trade Mark), and squalene in a ratio of 0.08: 1.0: 2.0

was employed.

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GnRH and anti-GnRH antibody titers were assayed by radioimmunosesay (RiA). Iodination of GnRH (5 ug) with 1 mc1 of carrier-free Na 125 1(Amersham) was carried out by the iodogen method (Braker PJ, Speck JC: Protein and cell

membrane iodination with a sparingly soluble chloramide 1,3,4,6-tetrachlroro-3,6 dippenylglycouril Blochem. Biophys. Res. Commun. 80: 849-855, 1978). Activity of 125_1 -labelled hormone ranged from 1,400-1,600 µci/µg.

The entibody titers, estimated in the assay system were expressed in terms of antigen-binding depacity [ABC]. All individual sera were titrated by dilution method simultaneously using the same batch of tracer. The assay protocol consisted of 50µl normal horse serum (diluted 2.5 times in assay buffer), 50µl of diluted 5.6 times in assay buffer), 50µl of diluted 5.6 times in assay buffer), 50µl of diluted 5.6 times in assay buffer (50µN, pH 7.4) and antiserum, 50 µl of phosphate buffer (50µN, pH 7.4) and 50µl of 125 1-LHRH, After incubation for 18 to 20 hours at 4°C, the antibody-bound fraction was separated by the method of Jeffcoate et al (Jeffcoate SL, Fraser HM, Holland DT, Gunn A: Radioimmnossasy of luteinizing hormone releasing bormone (LHRH) in serum from man, sheep and ret. Acts Endocrinol. (Copenh). 75:625-635, 1974).

Antigen-binding capacity (ng per ml) was calculated at a point at which proportionality between antiserum dilution point at which proportionality between antisecum dilution

Testosterone was determined by RIA, using labelled testosterone, with standards and antiserum to testosterone supplied by the World Health Organisation (WHO) under the matched Assay Reagents program.

.banistdo saw gnibnid HRH.1-1 2X1 bns

All the rats immunised with the conjugated vaccine developed antibodies against GRRH. With the rise in antibody titres, there occurred a concomitant fall in

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Synthesis of the vaccine was based on the premise that modification in the peptide backbone was mandatory for creating a defined site for conjugation with the carrier, without which a "self" hormone such as GnRH would not be immunogenic. Insertion of a D emino acid at from degradation (Monahan MW, Amoss MS, Anderson HA: Synthetic analogues of the hypothalamic luteintizing hormone releasing hormone with increased agonist or analogues. Biochemistry 12:4616-4620, 1973).

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An examination of tissues was effected ten weeks after immunisation. The data from such examination which is shown in Figure 2 of the drawings projects the marked drastic decrease in the prostate of the animals receiving the vaccine. The survival rate of the immunised animals the vaccine. The survival rate of the immunised animals and the vaccine. The survival rate of the immunised animals of the vaccine. The survival rate of the immunised animals and the vaccine. The survival rate of the immunised animals was virtually 100%. Anierior pituitary, adrenal and appear weights were not significantly altered after

male sex hormone levels as can be observed from Fig. 1 of the accompanying drawings which shows antigen binding capacity [ABC] and testosterone levels in rats immunized with the immunogenic substance. Each rat generated bioeffective antibodies of high titres showing the consistent immunogenicity of the preparation according to

so as to utilize its amino group for optional linkage to E-amino caproic acid, B-alanine or another non-protein amino acid. The results establish the fact that the modified GnRH analogue, conjugated to DT, produces an antibody response that is consistent and bioeffective.

The efficacy of the vaccine preparation for the difference or an analogue of the vaccine preparation for the difference or an analogue of the vaccine preparation.

been demonstrated (Sharpe RM, Fraser HM, Cooper I, not excluded. A local action of GnRH in the tests has deprivation of androgens, additional considerations are exercising the atrophic influence on prostate by Although it is likely that anti GnRH immunization is Eudocrinology 122:552-552). prostate after castration. yctivation of programmed cell death in the rat ventral spown by Kyprianou and Isaacs (Kyprianou N, Isaacs JT: castration-induced involution of the rat ventral prostate tmmunization on the prostate are also analogous to the The effects of rats, Endocrinol. 99:131-139, 1983). epithelial and stromal cells from immature and mature steroid dehydrogenase activities in ventral prostate Bird CE, Clark AF: androgen 5a reductase and 3a hydroxy metabolic activity of the prostate depends (Orlowski J, to be a definitive intracellar androgen upon which the it is converted to 5a dihydrotestosterone, now considered besses trom plasma to prostatic epithelial cells, where primarily dependent on androgenic stimuli. Testosterone Growth and function of the prostate are demonstrated. brognerud werked strobyl of the prostate was clearly

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Rommerts FFG: The secretion, messurement and function of testicular-LHRH like factor. Ann MY, Acad. Sci. 383:272-294, 1982). Whether or not GnRH exercises a direct action on the prostate is not known. Recently, however, Sheth et al. (Sheth AR, Joseph R, Maitra A: in vitro sheth et al. (Sheth AR, Joseph R, Maitra A: in vitro sheth et al. (Sheth AR, Joseph R, Maitra A: in vitro sheth et al. (Sheth AR, Joseph R, Maitra A: in vitro sheth et al. (Sheth AR, Joseph R, Maitra A: in vitro sheth et al. (Sheth AR, Joseph R, Maitra A: in vitro sheth et al. (Sheth AR, Joseph R, Maitra A: in vitro sheth et al. (Sheth AR, Joseph R, Maitra A: in vitro sheth et al. (Sheth AR, Joseph R, Maitra A: in vitro sheth et al. (Sheth AR, Joseph R, Maitra A: Josep

Although the exact mechanisms by which GnRH immunization interferes with prostatic growth and function need further clarification, it is obvious that their ability to inhibit gonadotropins, and consequently androgens, clearly parallels their delecterious effects.

Whilst in Example 4 the vaccine containing the peptide-protein conjugate has been specifically described in relation to its effect on the prostate the example

slso shows that the vaccine causes a marked reduction in

In this respect it will be appreciated by those skilled in the art that since GnRH is a master molecule controlling fertility in both male and female animals, this vaccine containing the peptide-immunogenic carrier protein conjugate or peptide-peptide dimer will be useful in all situations where an antagonist of LHRH may be usefully used, e.g. the control of male and female

weight of other reproductive organs.

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fertility, the suppression of heat in domestic pets, the suppression of heat in contraceptive. The intended to cover these other uses of the intended to cover these other uses of the peptide-immunogenic certies protein conjugate and

peptide-peptide dimer.

roup 3at Preimmunization Post immunization(9 weeks)					
energen pinding Anergen binding	festes sate	Testes size			
7400	05°T × 01°1	2.20 × 2.10	τ	TC-HRH	
099	02.1 × 00.1	2.15 × 2.10	ŧ	Satas	
0292	08.0 × 00.1	02.5 × 00.5	Ξ		
3230	02°T × 01°	2,35 × 2,20	÷		
0081	05*0 × 07*1	2.20 × 2.40	£		
7100	02°T × 06°	5°32 × 5°30	5		
0587	08.0 × 01.	2.50 x 2.40	÷		
7300	06.0 × 0	2°10 × 5°20	8		
7800	09.0 × 01.				
7400	05*0> × 08*	02.5 × 25.2	01		
5400	02.2 x 00.	2.40 x 2.40 Z	7	HDE EKH-DZ	
0087	01.1 × 06.	2°52 × 5°60 0	2		
7,80	08.0 × 06.	2.10 × 1.80	ε		
5400	02.0> x 0T.	0 02.2 × 01.2	,		
5897	02.1 × 0E.	3°20 × 3°40	s		
2000	08-0 × 02-	Z.30 × 2.20	9		
7800	02.0> x 08.		,		
5200	05.0> × 09.		8		
0251	08.0 × 01.		6		
1230	09.0 × 00.	2.30 × 2.20	OT		

TABLE I: Pre-and post immunization testes size and antibody titres of individual rate.

CITYIME:

.. A conjugate of the formula:

Y-orq-pra-lys-lys-lys-les-Arg-Pro-Y

wherein:

Pyr = pyroglutamic acid

His = histidine

Trp = tryptophan

Ser = serine

Tyr = tyrosine

D.Lys = D-lysine

ren = rencine

aginipas = pra

15 Arg = arginine

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bro = proline

A = GJA NHS OR NHEF

-qrT-ziH-ryq no nistoric cerrier protein or Pyr-His-Trp-

Ser-Tyr-D. Lys-Leu-Arg-Pro-Y as defined above.

2. An immunogenic substance capable of raising entibodies to GRRH in a mammalian subject, which immunogenic substance comprises a conjugate of claim 1.

25 3. A conjugate according to claim 1 wherein the immunogenic carrier protein is diphtheria toxoid (DT) or tetanus toxoid (TT).

y conjudate according to claim i or claim 3 wherein

the D-lysine residue is provided with a non-protein amino acid substituent to define the molar ratio between the peptide and protein.

- 5 5. A conjugate according to claim 4 wherein the nonprotein maino acid is selected from 6-aminocaproic acid
 or 8-alanine.
- 6. A conjugate according to any one of claims 1, 3, 4 $\,$ 10 or 5 which has been absorbed on alum or calcium
- 7. A conjugate according to anyone of claims 1, 3, 4, 5 or 6 for pharmaceutical use.

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phosphate.

- 8. A preparation comprising a conjugate according to any one of claims 1, 3, 4, 5 or 6 in combination with an adjuvant.
- 20 9. A preparation according to claim 7 wherein the adjuvant comprises nor-muramyl dipeptide or a sodium phthalylated derivative of Salmonella enteritidis lipopolysaccharide.
- 25 10. A method which comprises using a conjugate according to any one of claims 1, 3, 4, 5 or 6 to prepare a vaccine which is capable of stimulating the production of antibodies against GnRH.

I.1. A method for preparing a conjugate according to any one of claims 1, 3, 4, 5 or 6 which comprises using glutaraldehyde or 1-(3-dimethyl-amino-propyl)-3-ethyl

carbodifinide to couple X to
Pyr-His-Trp-Ser-Tyr-D.Lys-Leu-Arg-Pro-Y as defined above,
via the D-lysine residue.

12. A method according to claim 11 which comprises 10 providing the D-lysine residue with a non-protein amino

13. A method according to claim 12 wherein the non-protein amino acid substituent is ε -aminocaproic acid or β -alenine.

14. A method according to any one of claims 11, 12 or 13 wherein the immunogenic cerrier protein is diphtheria toxold (DT); tetanus toxoid (TT)

15. A conjugate substantially as described herein.

16. A method for preparing a conjugate substantially as described herein.

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ectd substituent.